# MULTITRACE -4- trace elements 4 injection, solution, concentrate American Regent, Inc.

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

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# MULTITRACE® - 4 CONCENTRATE (TRACE ELEMENTS INJECTION 4, USP)

FOR IV USE AFTER DILUTION

**Rx Only** 

## **DESCRIPTION**

**MULTITRACE® - 4 CONCENTRATE (TRACE ELEMENTS INJECTION 4, USP)** is a sterile nonpyrogenic solution containing four Trace Elements for use as an additive for Total Parenteral Nutrition (TPN).

Each mL provides: Zinc 5 mg, Copper 1 mg, Manganese 0.5 mg and Chromium 10 mcg.

Each mL contains: Zinc Sulfate Heptahydrate 22 mg (equivalent to 5 mg Zinc); Cupric Sulfate Pentahydrate 3.93 mg (equivalent to 1 mg Copper); Manganese Sulfate Monohydrate 1.54 mg (equivalent to 0.5 mg Manganese); Chromic Chloride Hexahydrate 51.3 mcg (equivalent to 10 mcg Chromium); and Water for Injection, q.s. pH of the solution may have been adjusted with Sulfuric Acid and/or Sodium Hydroxide. The 10 mL Multiple Dose Vial contains 0.9% Benzyl Alcohol as an antimicrobial preservative.

## CLINICAL PHARMACOLOGY

**ZINC** has been identified as a cofactor for over 70 different enzymes, including alkaline phosphatase, lactic dehydrogenase and both RNA and DNA polymerase. Zinc facilitates wound healing, helps maintain normal growth rates, normal skin hydration and senses of taste and smell.

Providing zinc during TPN prevents development of the following deficiency symptoms: parakeratosis, hypogeusia, anorexia, dysosmia, geophagia, hypogonadism, growth retardation and hepatosplenomegaly. At plasma levels below 20 mcg zinc/100 mL, dermatitis followed by alopecia has been reported for TPN patients.

**COPPER** is essential as a cofactor for serum ceruloplasmin, an oxidase necessary for proper formation of the iron carrier protein, transferrin. Copper also helps maintain normal rates of red and white blood cell formation. Scorbutic type bone changes seen in infants fed exclusively with copper-poor cow's milk are believed due to decreased activity of ascorbate oxidase, a cuproenzyme.

Providing copper during TPN prevents development of the following deficiency symptoms: leukopenia, neutropenia, anemia, depressed ceruloplasmin levels, impaired transferrin formation and secondary iron deficiency.

**MANGANESE** is an activator for enzymes such as polysaccharide polymerase, liver arginase, cholinesterase and pyruvate carboxylase.

Providing manganese during TPN prevents development of the following deficiency symptoms: nausea and vomiting, weight loss, dermatitis, and changes in growth and color of hair.

**CHROMIUM** (trivalent) is part of glucose tolerance factor, an activator of insulin-mediated reactions. Chromium helps to maintain normal glucose metabolism and peripheral nerve function.

Providing chromium during TPN prevents development of the following deficiency symptoms: impaired glucose tolerance, ataxia, peripheral neuropathy, and a confusional state similar to mild/moderate hepatic encephalopathy.

## INDICATIONS AND USAGE

This formulation is indicated for use as a supplement to intravenous solutions given for TPN. Administration of the solution in TPN solutions helps to maintain plasma levels of zinc, copper, manganese, and chromium and to prevent depletion of endogenous stores of these trace elements and subsequent deficiency symptoms.

## CONTRAINDICATIONS

**MULTITRACE**<sup>®</sup> - **4 CONCENTRATE** should not be given undiluted by direct injection into a peripheral vein because of the potential of infusion phlebitis.

#### **WARNINGS**

Copper and Manganese are eliminated via the bile. In patients with severe liver dysfunction and/or biliary tract obstruction, decreasing or omitting copper and manganese supplements entirely may be necessary.

This product contains aluminum that may be toxic. Aluminum may reach toxic levels with prolonged parenteral administration if kidney function is impaired. Premature neonates are particularly at risk because their kidneys are immature, and they require large amounts of calcium and phosphate solutions, which contain aluminum.

Research indicates that patients with impaired kidney function, including premature neonates, who receive parenteral levels of aluminum at greater than 4 to 5 mcg/kg/day accumulate aluminum at levels associated with central nervous system and bone toxicity. Tissue loading may occur at even lower rates of administration.

## **PRECAUTIONS**

Before administering MULTITRACE®- 4 CONCENTRATE in TPN solutions, the physician must assess the metabolic requirements for trace elements and disease state of the patient. Frequent determinations of serum levels of the various trace elements are suggested as a guideline for adjusting the dosage or completely omitting the solution. ZINC is eliminated via the intestine and kidneys. The possibility of retention should be considered in patients with malfunctioning excretory routes. COPPER and MANGANESE are eliminated via the bile, therefore, the possibility of the retention of these elements should be considered in patients with biliary obstruction. Ancillary routes of MANGANESE excretion, however, include pancreatic juice, or reabsorption into the lumen of duodenum, jejunum, or ileum.

In assessing the contribution of **CHROMIUM** supplements to maintenance of normal glucose homeostasis, consideration should be given to the possibility that the patient may be diabetic, in which case oral or intravenous antidiabetic medication may be indicated.

## **Pregnancy**

Teratogenic Effects

Pregnancy Category C: Safety for use in pregnancy has not been established. Use of **MULTITRACE**® - **4 CONCENTRATE** in women of childbearing potential requires that anticipated benefits be weighed against possible hazards.

## ADVERSE REACTIONS

The amounts of **ZINC**, **COPPER**, **MANGANESE**, AND **CHROMIUM** in the solution are very small and toxicity symptoms due to these trace elements at suggested dosage levels are considered unlikely to occur.

#### **OVERDOSAGE**

Symptoms of **ZINC** overdosage resulting from oral ingestion of Zinc Sulfate in large amounts have resulted in death. Symptoms included nausea, vomiting, dehydration, electrolyte imbalances, dizziness, abdominal pain, lethargy and incoordination. Single intravenous doses of 1 to 2 mg zinc/kg body weight have been given to adult leukemic patients without toxic manifestations. Normal plasma levels for Zinc vary from approximately 88 to 112 mcg/100 mL. Plasma levels sufficient to produce symptoms of toxic manifestations are not known. Calcium supplements may confer a protective effect against Zinc toxicity.

Symptoms of **COPPER** toxicity reported in literature include prostration, behavior change, diarrhea, progressive marasmus, hypotonia, photophobia and peripheral edema. D-penicillamine has been reported effective as an antidote.

**MANGANESE** toxicity has not been reported in patients receiving TPN. Neither have reports of manganese toxicity from excessive intake in foods and/or beverages been published.

Symptoms of **CHROMIUM** toxicity include nausea, vomiting, ulcers and gastrointestinal tract, renal and hepatic damage, and abnormalities of the central nervous system culminating in convulsions and coma. Trivalent Chromium administered intravenously to TPN patients has been shown to be nontoxic when given at dosage levels up to 250 mcg/day for two consecutive weeks.

## DOSAGE AND ADMINISTRATION

Each mL of the solution provides Zinc 5 mg, Copper 1 mg, Manganese 0.5 mg, and Chromium 10 mcg. The suggested dosage ranges for the four trace elements are:

#### ZINC

For the metabolically stable adult receiving TPN, the suggested intravenous dosage level is 2.5 to 4 mg zinc/day. An additional 2 mg zinc/day is suggested for acute catabolic states. For the stable adult with fluid loss from the small bowel, an additional 12.2 mg zinc/liter of small bowel fluid lost, or an additional 17.1 mg zinc/kg of stool or ileostomy output is recommended. Frequent monitoring of zinc blood levels is suggested for patients receiving more than the usual maintenance dosage level of zinc. Normal plasma levels for zinc vary from approximately 88 to 112 mcg/100 mL.

For full term infants and children up to 5 years of age, 100 mcg zinc/kg/day is recommended. For premature infants (birth weight less than 1500 g) up to 3 kg in body weight, 300 mcg zinc/kg/day is suggested.

#### COPPER

For the metabolically stable adult receiving TPN, the suggested additive dosage level is 0.5 to 1.5 mg copper/day. For pediatric patients, the suggested additive dosage level is 20 mcg copper/kg/day. The normal plasma range for copper is approximately 80 to 160 mcg/100 mL.

## **MANGANESE**

For the metabolically stable adult receiving TPN, the suggested additive dosage level for manganese is 0.15 to 0.8 mg/day. For pediatric patients, a dosage level of 2 to 10 mcg manganese/kg/day is recommended.

## **CHROMIUM**

For the metabolically stable adult receiving TPN, the suggested additive dosage level is 10 to 15 mcg chromium/day. The metabolically stable adult with intestinal fluid loss may require 20 mcg chromium/day with frequent monitoring of blood levels as a guideline for subsequent administration. For pediatric patients, the suggested additive dosage level is 0.14 to 0.20 mcg/kg/day.

Aseptic addition of  $MULTITRACE^{\$}$  - 4 CONCENTRATE to parenteral nutrition solutions under a laminar flow hood is recommended. The trace elements present in  $MULTITRACE^{\$}$  - 4 CONCENTRATE are physically compatible with the electrolytes and vitamins usually present in parenteral nutrition formulations.

Do not directly mix ascorbic acid injection with copper or selenium containing parenteral products in the same syringe or vial, as this admixture may cause the formation of an insoluble precipitate.

Parenteral drug products should be inspected visually for particulate matter and discoloration prior to administration, whenever solution and container permit.

Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) (See USP Controlled Room Temperature).

## **HOW SUPPLIED**

## MULTITRACE® - 4 CONCENTRATE (TRACE ELEMENTS INJECTION 4, USP)

Each mL provides: Zinc 5 mg, Copper 1 mg, Manganese 0.5 mg, and Chromium 10 mcg.

NDC 0517-7201-25 1 mL Single Dose Vial Packaged in boxes of 25

NDC 0517-7210-25 10 mL Multiple Dose Vial\* Packaged in boxes of 25

AMERICAN REGENT, INC. SHIRLEY, NY 11967

IN7201 Rev. 8/18

#### PRINCIPAL DISPLAY PANEL - 1 mL

**Container** 

NDC 0517-7201-25

MULTITRACE -4 CONCENTRATE (TRACE ELEMENTS INJECTION 4, USP)

1 mL SINGLE DOSE VIAL

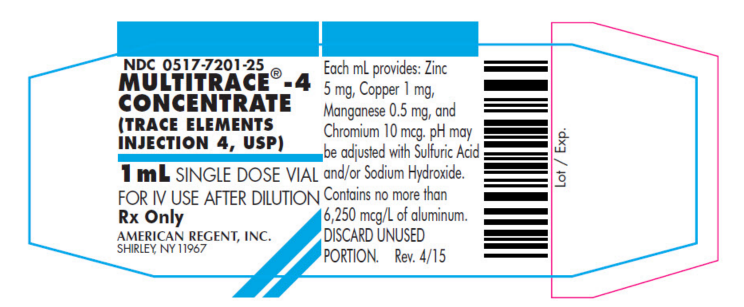
FOR IV USE AFTER DILUTION

**Rx Only** 

AMERICAN REGENT, INC.

SHIRLEY, NY 11967

<sup>\*</sup>Contains 0.9% Benzyl Alcohol as an antimicrobial preservative.



## Carton

MULTITRACE -4 CONCENTRATE (TRACE ELEMENTS INJECTION 4, USP)

NDC 0517-7201-25 25 x 1 mL SINGLE DOSE VIALS

## FOR IV USE AFTER DILUTION - PRESERVATIVE FREE

## **Rx Only**

Each mL provides: Zinc 5 mg, Copper 1 mg, Manganese 0.5 mg, and Chromium 10 mcg. Each mL contains: Zinc Sulfate (Heptahydrate) 22 mg, Cupric Sulfate (Pentahydrate) 3.93 mg, Manganese Sulfate (Monohydrate) 1.54 mg, Chromic Chloride (Hexahydrate) 51.3 mcg and Water for Injection q.s. pH may be adjusted with Sulfuric Acid and/or Sodium Hydroxide. Sterile, nonpyrogenic.

**WARNING**: DISCARD UNUSED PORTION – USE ONLY IF SOLUTION IS CLEAR. Store at 20°-25°C (68°-77°F); excursions permitted to 15°-30°C (59°-86°F) (See USP Controlled Room Temperature).

Directions for Use: See Package Insert.

AMERICAN REGENT, INC. SHIRLEY, NY 11967

Rev. 11/05

# MULTITRACE®-4 CONCENTRATE

(TRACE ELEMENTS INJECTION 4, USP)

NDC 0517-7201-25 25 x 1 mL SINGLE DOSE VIALS

## FOR IV USE AFTER DILUTION-PRESERVATIVE FREE

Rx Only

Exp.

Each mL provides: Zinc 5 mg, Copper 1 mg, Manganese 0.5 mg, and Chromium 10 mcg. Each mL contains: Zinc Sulfate (Heptahydrate) 22 mg, Cupric Sulfate (Pentahydrate) 3.93 mg, Manganese Sulfate (Monohydrate) 1.54 mg, Chromic Chloride (Hexahydrate) 51.3 mcg and Water for Injection q.s. pH may be adjusted with Sulfuric Acid and/or Sodium Hydroxide. Sterile, nonpyrogenic. WARNING: DISCARD UNUSED PORTION - USE ONLY IF SOLUTION IS CLEAR. Store at 20°-25°C (68°-77°F); excursions permitted to 15°-30°C (59°-86°F) (See USP Controlled Room Temperature).

Directions for Use: See Package Insert.

AMERICAN REGENT, INC. SHIRLEY, NY 11961



PRINCIPAL DISPLAY PANEL - 10 mL

Container

NDC 0517-7210-25

MULTITRACE -4 CONCENTRATE (TRACE ELEMENTS INJECTION 4, USP)

10 mL MULTIPLE DOSE VIAL FOR IV USE AFTER DILUTION Rx Only

AMERICAN REGENT, INC. SHIRLEY, NY 11967



MULTIPLE DOSE VIAL

FOR IV USE AFTER DILUTION

RX Only

(Hexanydraie) 51.3 mcg,
Benzyl Alcohol 0.9% as an
antimicrobial preservative and
Water for Injection q.s.
pH may be adjusted with

AMERICAN REGENT, INC. SHIRLEY, NY 11967 Each mL provides: Zinc 5 ma, Copper 1 mg, Manganese 0.5 mg, and Chromium 10 mcg. Each mL contains: Zinc Sulfate (Heptahydrate) 22 mg, Cupric Sulfate (Pentahydrate) 3.93 mg, Manganese Sulfate (Monohydrate) 1.54 mg, Chromic Chloride (Hexahydrate) 51.3 mcg, Benzyl Alcohol 0.9% as an Water for Injection q.s. pH may be adjusted with Sulfuric Acid and/or Sodium Hydroxide. Contains no more than 6,250 mcg/L of aluminum. Rev. 4/15



**Carton** 

MULTITRACE -4 CONCENTRATE (TRACE ELEMENTS INJECTION 4, USP)

NDC 0517-7210-25 25 x 10 mL MULTIPLE DOSE VIALS

FOR IV USE AFTER DILUTION

## **Rx Only**

Each mL provides: Zinc 5 mg, Copper 1 mg, Manganese 0.5 mg, Chromium 10 mcg. Each mL contains: Zinc Sulfate (Heptahydrate) 22 mg, Cupric Sulfate (Pentahydrate) 3.93 mg, Manganese Sulfate (Monohydrate) 1.54 mg, Chromic Chloride (Hexahydrate) 51.3 mcg, Benzyl Alcohol 0.9% as an antimicrobial preservative, and Water for Injection q.s. pH may be adjusted with Sulfuric Acid and/or Sodium Hydroxide. Sterile, nonpyrogenic.Store at 20°-25°C (68°-77°F); excursions permitted to 15°-30°C (59°-86°F) (See USP Controlled Room Temperature).

Directions for Use: See Package Insert. **AMERICAN** 

**REGENT, INC.** SHIRLEY, NY 11967

Rev. 11/05

# MULTITRACE®-4 CONCENTRATE

(TRACE ELEMENTS INJECTION 4, USP)

NDC 0517-7210-25 25 x 10 mL MULTIPLE DOSE VIALS

#### FOR IV USE AFTER DILUTION

Rx Only

Each mL provides: Zinc 5 mg, Copper 1 mg, Manganese 0.5 mg, Chromium 10 mcg. Each mL contains: Zinc Sulfate (Heptahydrate) 22 mg, Cupric Sulfate (Pentahydrate) 3.93 mg, Manganese Sulfate (Monohydrate) 1.54 mg, Chromic Chloride (Hexahydrate) 51.3 mcg, Benzyl Alcohol 0.9% as an antimicrobial preservative, and Water for Injection q.s. pH may be adjusted with Sulfuric Acid and/or Sodium Hydroxide. Sterile, nonpyrogenic.

Store at 20°-25°C (68°-77°F); excursions permitted to 15°-30°C (59°-86°F) (See USP Controlled

Room Temperature).

Directions for Use: See Package Insert.

AMERICAN REGENT, INC. SHIRLEY, NY 11967

Rev. 11/05



## Serialization Label - 1 mL





## **MULTITRACE -4**

trace elements 4 injection, solution, concentrate

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0517-7201
Route of Administration	INTRAVENOUS		

## **Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
ZINC SULFATE HEPTAHYDRATE (UNII: N57JI2K7WP) (ZINC CATION - UNII: 13S 1S8 SF37)	ZINC CATION	22 mg in 1 mL
CUPRIC SULFATE (UNII: LRX7AJ16DT) (CUPRIC CATION - UNII:8CBV67279L)	CUPRIC CATION	3.93 mg in 1 mL
MANGANESE SULFATE (UNII: W00LYS4T26) (MANGANESE CATION (2+) - UNII:H6 EP7W5457)	MANGANESE CATION (2+)	1.54 mg in 1 mL
CHROMIC CHLORIDE (UNII: KB1PCR9 DMW) (CHROMIC CATION - UNII:X1N450 8 KF1)	CHROMIC CATION	51.3 ug in 1 mL

Inactive Ingredients		
Ingredient Name	Strength	
SULFURIC ACID (UNII: O40 UQP6 WCF)		
SO DIUM HYDRO XIDE (UNII: 55X04QC32I)		
WATER (UNII: 059QF0KO0R)		

]	Packaging			
7	t Item Code	Package Description  Marketing Start Date		Marketing End Date
1	NDC:0517-7201- 25	25 in 1 TRAY	12/07/1993	
1	L	1 mL in 1 VIAL, SINGLE-DOSE; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
UNAPPROVED DRUG OTHER		12/07/1993	

## **MULTITRACE -4**

trace elements 4 injection, solution, concentrate

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0517-7210
Route of Administration	INTRAVENOUS		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
ZINC SULFATE HEPTAHYDRATE (UNII: N57JI2K7WP) (ZINC CATION - UNII: 13S1S8SF37)	ZINC CATION	22 mg in 1 mL	
CUPRIC SULFATE (UNII: LRX7AJ16DT) (CUPRIC CATION - UNII:8CBV67279L)	CUPRIC CATION	3.93 mg in 1 mL	
MANGANESE SULFATE (UNII: W00LYS4T26) (MANGANESE CATION (2+) - UNII:H6EP7W5457)	MANGANESE CATION (2+)	1.54 mg in 1 mL	
CHROMIC CHLORIDE (UNII: KB1PCR9 DMW) (CHROMIC CATION - UNII:X1N450 8 KF1)	CHROMIC CATION	51.3 ug in 1 mI	
CIMO INC. CIMO INC. (CIMO INC. CIMO	GIRCOINE GITTOIT	51.5 ug m 11	

Inactive Ingredients				
Ingredient Name	Strength			
BENZYL ALCOHOL (UNII: LKG8494WBH)				
SULFURIC ACID (UNII: O40 UQP6 WCF)				
SO DIUM HYDRO XIDE (UNII: 55X04QC32I)				
WATER (UNII: 059QF0KO0R)				

I	Packaging				
#	Item Code	Package Description  Marketing Start Date		Marketing End Date	
1	NDC:0517-7210- 25	25 in 1 TRAY	11/29/1993		
1		10 mL in 1 VIAL, MULTI-DOSE; Type 0: Not a Combination Product			

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
UNAPPROVED DRUG OTHER		11/29/1993	

## Labeler - American Regent, Inc. (002033710)

Establishment			
Name	Address	ID/FEI	Business Operations
American Regent, Inc.		002033710	ANALYSIS(0517-7201, 0517-7210), MANUFACTURE(0517-7201, 0517-7210), STERILIZE(0517-7201, 0517-7210)

Revised: 4/2019 American Regent, Inc.